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NEWS 1 Web Page for STN Seminar Schedule - N. America
NEWS 2 JUL 02 LMEDLINE coverage updated
NEWS 3 JUL 02 SCISEARCH enhanced with complete author names
NEWS 4 JUL 02 CHEMCATS accession numbers revised
NEWS 5 JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS 6 JUL 16 CAplus enhanced with French and German abstracts
NEWS 7 JUL 18 CA/CAplus patent coverage enhanced
NEWS 8 JUL 26 USPATFULL/USPAT2 enhanced with IPC reclassification
NEWS 9 JUL 30 USGENE now available on STN
NEWS 10 AUG 06 CAS REGISTRY enhanced with new experimental property tags
NEWS 11 AUG 06 FSTA enhanced with new thesaurus edition
NEWS 12 AUG 13 CA/CAplus enhanced with additional kind codes for granted patents
NEWS 13 AUG 20 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 14 AUG 27 Full-text patent databases enhanced with predefined patent family display formats from INPADOCDB
NEWS 15 AUG 27 USPATOLD now available on STN
NEWS 16 AUG 28 CAS REGISTRY enhanced with additional experimental spectral property data
NEWS 17 SEP 07 STN AnaVist, Version 2.0, now available with Derwent World Patents Index
NEWS 18 SEP 13 FORIS renamed to SOFIS
NEWS 19 SEP 13 INPADOCDB enhanced with monthly SDI frequency
NEWS 20 SEP 17 CA/CAplus enhanced with printed CA page images from 1967-1998
NEWS 21 SEP 17 CAplus coverage extended to include traditional medicine patents
NEWS 22 SEP 24 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 23 OCT 02 CA/CAplus enhanced with pre-1907 records from Chemisches Zentralblatt
NEWS 24 OCT 19 BEILSTEIN updated with new compounds
NEWS 25 NOV 15 Derwent Indian patent publication number format enhanced
NEWS 26 NOV 19 WPIX enhanced with XML display format
NEWS 27 NOV 30 ICSD reloaded with enhancements
NEWS 28 DEC 04 LINPADOCDB now available on STN

NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

NEWS HOURS STN Operating Hours Plus Help Desk Availability
NEWS LOGIN Welcome Banner and News Items
NEWS IPC8 For general information regarding STN implementation of IPC 8

Page 2

Enter NEWS followed by the item number or name to see news on that specific topic.

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STRUCTURE FILE UPDATES: 7 DEC 2007 HIGHEST RN 957187-88-1
DICTIONARY FILE UPDATES: 7 DEC 2007 HIGHEST RN 957187-88-1

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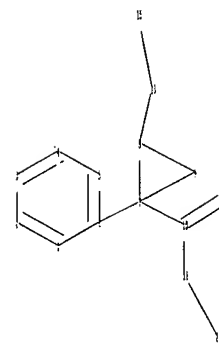
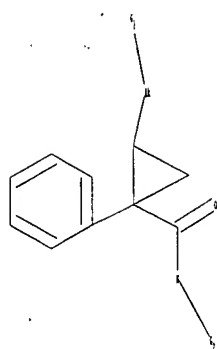
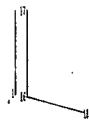
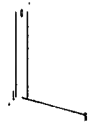
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Uploading C:\Program Files\Stnexp\Queries\10541047\Struc 3.str



chain nodes :

10 11 12 13 15 16 17 18 21

ring nodes :

1 2 3 4 5 6 7 8 9

chain bonds :

6-7 7-10 8-13 10-11 10-12 11-21 13-15 16-17 16-18

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-9 8-9

exact/norm bonds :

7-8 7-9 8-9 8-13 10-11 10-12 11-21 13-15 16-17 16-18

exact bonds :

6-7 7-10

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

G1:H,X

G2:Hy, [*1]

10541047.trn

Match level :
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS
11:CLASS 12:CLASS 13:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 21:CLASS

L1 STRUCTURE UPLOADED

=> d
L1 HAS NO ANSWERS
L1 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY -- AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

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SAMPLE SCREEN SEARCH COMPLETED - 906 TO ITERATE

100.0% PROCESSED 906 ITERATIONS 0 ANSWERS
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 16315 TO 19925
PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> l1 full
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FULL SCREEN SEARCH COMPLETED - 18486 TO ITERATE

100.0% PROCESSED 18486 ITERATIONS 9 ANSWERS
SEARCH TIME: 00.00.01

L3 9 SEA SSS FUL L1

=> file caplus
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FULL ESTIMATED COST ENTRY SESSION
172.10 172.31

FILE 'CAPLUS' ENTERED AT 17:55:42 ON 09 DEC 2007
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FILE COVERS 1907 - 9 Dec 2007 VOL 147 ISS 25
FILE LAST UPDATED: 7 Dec 2007 (20071207/ED)

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They are available for your review at:

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=> l3
L4 3 L3

=> d ibib abs hitstr 1-3

L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2007:1176083 CAPLUS

DOCUMENT NUMBER: 147:469226

TITLE: Indolyl cycloalkylcarboxamide compounds as modulators of ATP-binding cassette transporters and their preparation, pharmaceutical compositions and use in the treatment of diseases

INVENTOR(S): Ruah, Sara S.; Hadida; Grootenhuis, Peter D. J.; Van Goor, Frederick; Zhou, Jinglan; Bear, Brian; Miller, Mark T.; McCartney, Jason; Numa, Mehdi Michel Jamel

PATENT ASSIGNEE(S): Vertex Pharmaceuticals Incorporated, USA

SOURCE: PCT Int. Appl., 276pp.

CODEN: PIKXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2007117715	A2	20071018	WO 2007-US8975	20070409
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MX, MY, NZ, NA, NG, NI, NO, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				

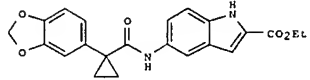
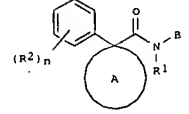
US 2007244159 A1 20071018 US 2007-786001 20070409

PRIORITY APPLN. INFO.: US 2006-790459 P 20060407

OTHER SOURCE(S): MARPAT 147:469226

GI

L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)



AB The invention relates to compds. of formula I and pharmaceutically acceptable compns. thereof, which are useful as modulators of ATP-Binding Cassette ("ABC") transporters or fragments thereof, including Cystic Fibrosis Transmembrane Conductance Regulator ("CFTR"). The invention also relates to methods of treating ABC transporter mediated diseases using compds. of the present invention. Compds. of formula I wherein R1 and R2 are independently H, (un)substituted (un)branched C1-6 aliphatic chain, halo, OH, NH2, NO2, CN, OCF3, etc.; Ring A is (un)substituted 3- to 7-membered mono(heterocyclic) ring; B is (un)substituted indolyl; n is 1-3; and their pharmaceutically acceptable salts thereof, are claimed. Example compound II

was prepared by chlorination of 1-(3,4-methylenedioxophenyl)cyclopropanecarbonylic acid followed by amidation with 5-aminoindole-2-carboxylic acid Et ester. All the invention compds. were evaluated for their ATP-binding cassette transporter modulatory activity. From the assay, it was determined

that the invention compds. exhibited EC50 values from about 3.8 nM to about 13.5 μM and the efficacies was found to be from about 35 % to about 110 %.

IT 952664-12-9 952664-38-9
RL: PAC (Pharmacological activity); SPA (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of indolyl cycloalkylcarboxamide compds.)

as ATP-binding cassette transporters useful in treatment of ABC transporter-mediated diseases

RN 952664-12-9 CAPLUS

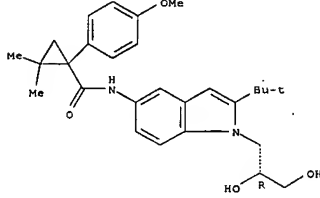
CH Cyclopropanecarboxamide, N-[1-[(2R)-2,3-dihydroxypropyl]-2-(1,1-dimethylethyl)-1H-indol-5-yl]-1-(4-methoxyphenyl)-2,2-dimethyl- (CA INDEX NAME)

INDEX

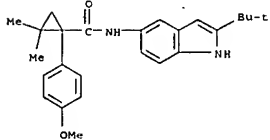
L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

NAME)

Absolute stereochemistry.



RN 952664-38-9 CAPLUS
CN Cyclopropanecarboxamide, N-[2-(1,1-dimethylethyl)-1H-indol-5-yl]-1-(4-methoxyphenyl)-2,2-dimethyl- (CA INDEX NAME)



L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:606457 CAPLUS

DOCUMENT NUMBER: 141:157108

TITLE: Preparation of aryl substituted cyclopropanecarboxamides for therapeutic use as glucokinase activators

INVENTOR(S): Weichert, Andreas Gerhard; Barrett, David Gene; Heuser, Stefan; Riedl, Rainer; Tebbe, Mark Joseph; Zaliani, Andrea

PATENT ASSIGNEE(S): Eli Lilly and Company, USA

SOURCE: PCT Int. Appl., 141 pp.

CODEN: PIKXD2

DOCUMENT TYPE: Patent

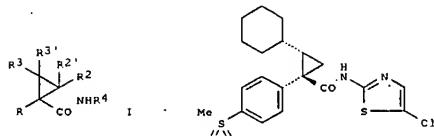
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004063179	A1	20040729	WO 2003-US37088	20031216
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RN 2509086	A1	20040729	CA 2003-2509086	20031216
AU 2003297291	A1	20040810	AU 2003-297291	20031216
EP 1585739	A1	20051019	EP 2003-815189	20031216
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JP 2006515858	T	20060608	JP 2004-566494	20031216
US 2006111353	A1	20060525	US 2005-541047	20050629

PRIORITY APPLN. INFO.: WO 2003-US37088 W 20031216

OTHER SOURCE(S): MARPAT 141:157108
GI

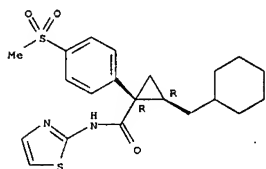
L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

AB Cyclopropylcarboxamides, such as I [R = substituted aryl or heteroaryl; R₂, R_{2'} = H, Me, halogen; R₃ = alkyl, cycloalkyl, cycloalkylmethyl, etc.; R_{3'} = H, halogen, alkyl, perfluoroalkyl; R₄ = heteroaryl, such as thiazolyl], were prepared for use in pharmaceutical compns. as glucokinase activators which are useful for treatment of type II diabetes. Thus, trans-cyclopropylcarboxamide II was prepared via an amidation reaction of the corresponding cyclopropanecarboxylic acid with (5-chlorothiazol-2-yl)amine hydrochloride using TBTU and Et₃N in THF. The prepared cyclopropylcarboxamides were assayed for their ability to increase glucokinase activity. Also, pharmaceutical formulations containing the prepared cyclopropylcarboxamides were presented.

IT 731016-85-6P
731016-90-3P 731019-01-5P 731019-21-9P
RN: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOl (Biological study); PREP (Preparation); USES (Uses)
(preparation of substituted aryl substituted cyclopropylcarboxamides for therapeutic use as glucokinase activators)

RN 731016-72-1 CAPLUS
CN Cyclopropanecarboxamide,
2-(cyclohexylmethyl)-1-[4-(methylsulfonyl)phenyl]-
N-2-thiazolyl-, (1R,2R)-rel- (CA INDEX NAME)

Relative stereochemistry.



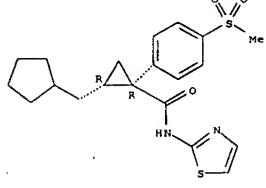
RN 731016-79-8 CAPLUS
CN Cyclopropanecarboxamide,
2-(2-methylpropyl)-1-[4-(methylsulfonyl)phenyl]-N-
2-thiazolyl-, (1R,2R)-rel- (CA INDEX NAME)

Relative stereochemistry.

L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

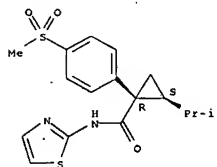
RN 731019-01-5 CAPLUS
CN Cyclopropanecarboxamide, 2-(cyclopentylmethyl)-1-[4-(methylsulfonyl)phenyl]-N-2-thiazolyl-, (1R,2R)-rel- (CA INDEX NAME)

Relative stereochemistry.

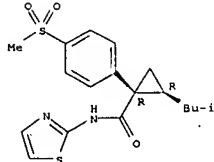


RN 731019-21-9 CAPLUS
CN Cyclopropanecarboxamide,
2-(1-methylethyl)-1-[4-(methylsulfonyl)phenyl]-N-
2-thiazolyl-, (1R,2S)-rel- (CA INDEX NAME)

Relative stereochemistry.

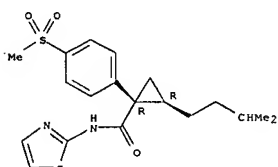


L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)



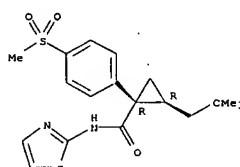
RN 731016-85-6 CAPLUS
CN Cyclopropanecarboxamide,
2-(3-pentyl)-1-[4-(methylsulfonyl)phenyl]-N-
2-thiazolyl-, (1R,2R)-rel- (CA INDEX NAME)

Relative stereochemistry.



RN 731016-90-3 CAPLUS
CN Cyclopropanecarboxamide, 2-(2,2-dimethylpropyl)-1-[4-(methylsulfonyl)phenyl]-N-2-thiazolyl-, (1R,2R)-rel- (CA INDEX NAME)

Relative stereochemistry.

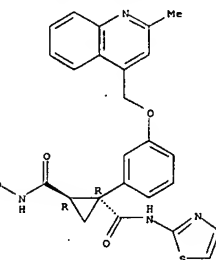
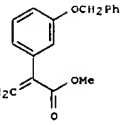
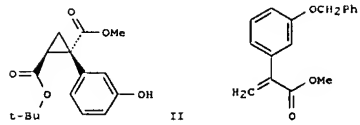
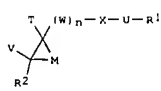


L4 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003511283 CAPLUS
DOCUMENT NUMBER: 139-85018
TITLE: Preparation of TNF- α -inhibiting hydroxamic or carboxylic acid functionalized cycloalkanes for the treatment of inflammatory disorders
INVENTOR(S): Zhu, Zhenming; Mazzola, Robert, Jr.; Guo, Zhuyan; Lavey, Brian J.; Sinning, Lisa; Kozlowski, Joseph; McKittrick, Brian; Shih, Neng-Yang
PATENT ASSIGNEE(S): Schering Corporation, USA
SOURCE: PCT Int. Appl.; 179 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003053915	A2	20030703	WO 2002-US40453	20021219
WO 2003053915	A3	20030918		
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CA 2470620	A1	20030703	CA 2002-2470620	20021219
AU 2002357885	A1	20030709	AU 2002-357885	20021219
US 2004038941	A1	20040226	US 2002-323511	20021219
US 6938466	B2	20050510		
EP 1458676	A2	20040922	EP 2002-792459	20021219
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HU 200500016	A2	20050428	HU 2005-16	20021219
JP 2005513125	T	20050512	JP 2005-554632	20021219
CN 1620424	A	20050525	CN 2002-828170	20021219
US 2004102418	A1	20040527	US 2003-716890	20031119
US 7034057	B2	20060425		
ZA 2004004586	A	20050823	ZA 2004-4586	20040609
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			US 2002-323511	A3 20021219
			WO 2002-US40453	W 20021219
			US 2003-716890	A3 20031119

OTHER SOURCE(S): MARPAT 139:85038
GI



AB This invention relates to compds. of formula I [M = -(C(R30)(R40))^m-, wherein m = 1-6; T = substituted alkyl, (un)substituted-cycloalkyl, -heterocycloalkyl, -aryl, etc.; V = (un)substituted alkyl, cycloalkyl, heteroaryl, etc.; R1 = (un)substituted alkyl, alkyne, alkene, cycloalkyl, aryl, etc.; R2 = H, halo, (un)substituted alkyl, cycloalkyl, etc.; U = bond, alkyl, heteroalkyl, heteroatoms; X = (un)substituted alkylene, cycloalkylene, arylene, etc.; W = carboxy, substituted iminomethylene, SO₂, SO, etc., wherein n = 0-2; R30 and R40 independently = H or halo,

CN, NO₂, (un)substituted alkyl, etc.; or R30 and R40 may be taken together with the atom to which they are attached to form C=O, with provisions] or a pharmaceutically acceptable salt, solvate or isomer thereof, which can be useful for the treatment of diseases or conditions mediated by MMPs, TNF-alpha or combinations thereof. Thus, II was prepared from Me methoxypheophenone with the cyclopropane ring diastereoselectively formed by cyclization of intermediate III with S-carbo-tert-butoxymethyltetrahydrophosphine bromide with subsequent hydrogenation and resolution of enantiomers. Numerous compds. of the invention possessed

K1 values of less than 20 nM in a TNF- α convertases (TACE) inhibitory activity assay. As TNF- α inhibitors, I will be useful in treatment of inflammatory disorders.

IT 556108-71-5
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of hydroxamic or carboxylic acid functionalized cycloalkanes as inhibitors of tumor necrosis factor alpha and/or matrix metalloproteinases)

RN 556108-71-5 CAPLUS

CN 1,2-Cyclop propane dicarboxamide, N2-hydroxy-1-[3-[(2-methyl-4-quinolinyl)methoxy]phenyl]-N1-2-thiazolyl-, (1R,2R)- (CA INDEX NAME)

Page 9

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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.34	-2.34

STN INTERNATIONAL LOGOFF AT 17:56:40 ON 09 DEC 2007

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